



The Influence of Genotype and Environment on Small RNA Profiles in Grapevine Berry

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Understanding the molecular mechanisms involved in the interaction between the genetic composition and the environment is crucial for modern viticulture. We approached this issue by focusing on the small RNA transcriptome in grapevine berries of the two varieties Cabernet Sauvignon and Sangiovese, growing in adjacent vineyards in three different environments. Four different developmental stages were studied and a total of 48 libraries of small RNAs were produced and sequenced. Using a proximity-based pipeline, we determined the general landscape of small RNAs accumulation in grapevine berries. We also investigated the presence of known and novel miRNAs and analyzed their accumulation profile. The results showed that the distribution of small RNA-producing loci is variable between the two cultivars, and that the level of variation depends on the vineyard. Differently, the profile of miRNA accumulation mainly depends on the developmental stage. The vineyard in Riccione maximizes the differences between the varieties, promoting the production of more than 1000 specific small RNA loci and modulating their expression depending on the cultivar and the maturation stage. In total, 89 known vvi-miRNAs and 33 novel vvi-miRNA candidates were identified in our samples, many of them showing the accumulation profile modulated by at least one of the factors studied. The *in silico* prediction of miRNA targets suggests their involvement in berry development and in secondary metabolites accumulation such as anthocyanins and polyphenols.

Keywords: Vitis vinifera, Genotype x Environment (GxE), small RNAs, miRNAs, high throughput sequencing, berry

INTRODUCTION

The ability of a genotype to produce different phenotypes as a function of environmental cues is known as phenotypic plasticity (Bradshaw, 1965; Sultan, 2000; Pigliucci, 2001; Gratani, 2014). Phenotypic plasticity is considered one of the main processes by which plants, as sessile organisms, can face and adapt to the spatio-temporal variation of environmental factors (Nicotra et al., 2010; Palmer et al., 2012; Gratani, 2014).

Grapevine (*Vitis vinifera* L.) berries are characterized by high phenotypic plasticity (Dal Santo et al., 2013) and a genotype (cultivar or clone) can present variability within berries, among berries

OPEN ACCESS

Edited by:

Mahmoud W. Yaish, Sultan Qaboos University, Oman

Reviewed by:

Nataliya V. Melnikova, Engelhardt Institute of Molecular Biology (RAS), Russia Douglas S. Domingues, Sao Paulo State University, Brazil

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Specialty section:

This article was submitted to Plant Physiology, a section of the journal Frontiers in Plant Science

Received: 19 February 2016 Accepted: 13 September 2016 Published: 05 October 2016

Citation:

Paim Pinto DL, Brancadoro L, Dal Santo S, De Lorenzis G, Pezzotti M, Meyers BC, Pè ME and Mica E (2016) The Influence of Genotype and Environment on Small RNA Profiles in Grapevine Berry. Front. Plant Sci. 7:1459. doi: 10.3389/fpls.2016.01459

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in a cluster, and among vines (Gray, 2002; Keller, 2010). Berry phenotypic traits, such as the content of sugars, acids, phenolic, anthocyanins, and flavor compounds, are the result of cultivar (G) and environmental influences (E), and often strong $G \times E$ interactions (Sadras et al., 2007). Although grapevine plasticity in response to environmental conditions and viticulture practices may provide advantages related to the adaptation of a cultivar to specific growing conditions, it may also cause irregular ripening (Selvaraj et al., 1994) and large inter-seasonal fluctuations (Clingeleffer, 2010), which are undesirable characteristics for wine making (Keller, 2010).

Due to its complex nature, the study of phenotypic plasticity is challenging and the mechanisms by which the genes affecting plastic responses operate are poorly characterized (Holloway, 2002; DeWitt and Scheiner, 2003; Nicotra et al., 2010; Gianoli and Valladares, 2012; Gratani, 2014). In fact it is often difficult to assess the performance of different phenotypes in different environments (Schmitt, 1993; Schmitt et al., 1999; Callaway et al., 2003).

It has been suggested that genetic and epigenetic regulation of gene expression might be at the basis of phenotypic plasticity through the activation of alternative gene pathways (Schlichting and Pigliucci, 1993; Pigllucci, 1996) or multiple genes (Lind et al., 2015). Epigenetics has been proposed as crucial in shaping plant phenotypic plasticity, putatively explaining the rapid and reversible alterations in gene expression in response to environmental changes. This fine-tuning of gene expression can be achieved through DNA methylation, histone modifications and chromatin remodeling (Goldberg et al., 2007; Geng et al., 2013; Duncan et al., 2014).

Small non-coding RNAs (small ncRNAs) are ubiquitous and adjustable repressors of gene expression across a broad group of eukaryotic species and are directly involved in controlling, in a sequence specific manner, multiple epigenetic phenomena such as RNA-directed DNA methylation and chromatin remodeling (Bernstein and Allis, 2005; Fagegaltier et al., 2009; Ha et al., 2009; Swami, 2010; Burkhart et al., 2011; Castel and Martienssen, 2013; Duncan et al., 2014) and might play a role in genotype by environment (GxE) interactions. In plants, small ncRNAs are typically 20-24 nt long RNA molecules and participate in a wide series of biological processes controlling gene expression via transcriptional and post-transcriptional regulation (Finnegan and Matzke, 2003; Kim, 2005; Chen, 2009; Guleria et al., 2011; Lelandais-Briere et al., 2012; Matsui et al., 2013). Moreover, small RNAs have been recently shown to play an important role in plants environmental plasticity (Formey et al., 2014; Borges and Martienssen, 2016).

Fruit maturation, the process that starts with fruit-set and ends with fruit ripening (Coombe, 1976), has been largely investigated in fleshy fruits such as tomato and grapevine. These studies highlighted, among others, the vast transcriptomic reprogramming underlying the berry ripening process (Guillaumie et al., 2011; Matas et al., 2011; Lijavetzky et al., 2012), the extensive plasticity of berry maturation in the context of a changing environment (Dal Santo et al., 2013; Gapper et al., 2014), and the epigenetic regulatory network which contributes to adjust gene expression to internal and external stimuli (Zhong et al., 2013; Liu et al., 2015). In particular, small RNAs, and especially microRNAs (miRNA), are involved, among others, in those biological processes governing fruit ripening (Karlova et al., 2013; Kullan et al., 2015).

In this work, we assessed the role of small ncRNAs in the plasticity of grapevine berries development, by employing next-generation sequencing. We focused on two cultivars of Vitis vinifera, Cabernet Sauvignon, and Sangiovese, collecting berries at four different developmental stages in three Italian vineyards, diversely located. First, we described the general landscape of small RNAs originated from hotspots present along the genome, examining their accumulation according to cultivars, environments and developmental stages. Subsequently, we analyzed miRNAs, identifying known and novel miRNA candidates and their distribution profiles in the various samples. Based on the in silico prediction of their targets, we suggest the potential involvement of this class of small RNAs in GxE interactions. The results obtained provide insights into the complex molecular machinery that connects the genotype and the environment.

MATERIALS AND METHODS

Plant Material

Two *V. vinifera* varieties Sangiovese (SG), a red Italian grape variety, and Cabernet Sauvignon (CS), an international variety, were grown side by side in three different Italian locations, representing traditional areas of Sangiovese cultivation in Italy with a long-standing winemaking tradition.

In order to reduce factors of variation, the age of the plants (between 10 and 12 years old), the clone type (Sangiovese clone R5 and Cabernet Sauvignon clone VCR23), the rootstock (*Vitis berlandieri* × *Vitis riparia*), the cultivation techniques (training system: low cordon; planting space: 2.40×0.8 m) and the health status were the same among all the locations.

The vineyards were located in Bolgheri (Bol), a coastal area of Tuscany, 50 m asl (above sea level) [GPS coordinates: SG 43.194090, 10.625186, CS 43.194622, 10.624392], in Montalcino (Mont) a mountain area of Tuscany, 195 m asl; [GPS coordinates: SG 42.980669, 11.433708, CS 42.985091, 11.435853] and in Riccione (Ric), a plain area of Emilia Romagna, 111 m asl; [GPS coordinates: SG 43.945261, 12.647235, CS 43.944372, 12.648995]. Further details on the environmental conditions of the vineyards are provided in Supplementary Figure 1.

Berries from four developmental stages were collected in two biological replicates, during the 2011 growing season, for a total of 48 samples (**Table 1**). The four sampled stages corresponded to pea size (ps), representing the first stage of berry development in this experimental plan, bunch closure (bc) also known as Lag Phase, 19-20 °Brix (19), which corresponds to 50% of sugar accumulation in berries, and harvest (hv), when the berries are fully ripened and the onset of sugar accumulation is over. About 200 berries per each developmental berry stage were sampled from upper, central and lower part of cluster, both from sunexposed and shaded side and split in two biological replicates. Per each vineyard, the berries were collected from about 20 vines selected in a single uniform row and immediately frozen in liquid nitrogen and stored at $-80^\circ \rm C$ prior to analysis.

The libraries were named using the initials of the vineyard where the berries were collected, followed by the initial of the cultivar and the developmental stage. For example, the sample containing berries of Sangiovese, collected in Montalcino at pea size, was named Mont_SG_ps.

RNA Extraction and Small RNA Libraries Construction

RNA extraction was performed as described in Kullan et al. (2015). Briefly, total RNA was extracted from 200 mg of ground berries pericarp tissue (entire berries without seeds) using 1 ml of Plant RNA Isolation Reagent (Life Technologies) following manufacturer's recommendations.

The small RNA fraction was isolated from the total RNA using the mirPremier[®] microRNA Isolation kit (Sigma-Aldrich) and dissolved in DEPC water. All the steps suggested in the technical bulletin for small RNA isolation of plant tissues were followed except the "Filter Lysate" step, which was omitted. The quality and quantity of small RNAs were evaluated by a NanoDrop 1000 spectrometer (Thermo Fisher Scientific), and their integrity assessed by an Agilent 2100 Bioanalyzer using a small RNA chip (Agilent Technologies) according to the manufacturer's instructions.

Small RNA libraries were prepared using the TruSeq Small RNA Sample Preparation Kit (Illumina[®]), following all manufacturers' instructions. Forty-eight bar-coded small RNA libraries were constructed starting from 50 ng of small RNAs. The quality of each library was assessed using an Agilent DNA 1000 chip for the Agilent 2100 Bioanalyzer. Libraries were grouped in pools with six libraries each (6-plex).

The pools of libraries were sequenced on an Illumina Hiseq 2000 at IGA Technology Services (Udine, Italy).

The sequencing data were submitted to GEO–NCBI under the accession number GSE85611.

Bioinformatics Analysis of Sequencing Data

Adaptor sequences were trimmed and only reads ranging from 18 to 34 nt in length after adapter removal were kept. Retained reads were mapped to the reference *Vitis vinifera* L. genomic sequence V1 (PN40024, Jaillon et al., 2007) using Bowtie (Langmead et al., 2009) and reads perfectly aligned to the genome were retained. Reads matching rRNAs, tRNAs, snRNAs, and snoRNAs were excluded.

Read counts were normalized by the linear count scaling method TP4M (transcripts per 4 million), in order to reduce sequencing bias and to allow the comparison of small RNA accumulation from different libraries. The normalized abundance was calculated as:

$$abundance = \left[\frac{raw \ value}{(total \ genome \ matches - t/r/sn/snoRNA / chloroplast \ mitochondria \ matches)} \right] \\ \times n_base$$

TABLE 1 | List of berry samples of *Vitis vinifera* used for the construction of the small RNA libraries.

Vineyard	Variety	Developmental Stages	Replicate	Library Code
	Cabernet	Pea size	1	Mont_CS_ps_
Montalcino	Sauvignon		2	Mont_CS_ps_
		Bunch closure	1	Mont_CS_bc_
			2	Mont_CS_bc_
		19 °Brix	1	Mont_CS_19_
			2	Mont_CS_19_
		Harvest	1	Mont_CS_hv_
			2	Mont_CS_hv_
Montalcino	Sangiovese	Pea size	1	Mont_SG_ps_
			2	Mont_SG_ps_
		Bunch closure	1	Mont_SG_bc_
			2	Mont_SG_bc_
		19 °Brix	1	Mont_SG_19_
			2	Mont_SG_19_
		Harvest	1	Mont_SG_hv_
			2	Mont_SG_hv_
Bolgheri	Cabernet Sauvignon	Pea size	1	Bol_CS_ps_1
Doighen	Sauvignon		2	Bol_CS_ps_2
		Bunch closure	1	Bol_CS_bc_1
			2	Bol_CS_bc_2
		19 °Brix	1	Bol_CS_19_1
			2	Bol_CS_19_2
		Harvest	1	Bol_CS_hv_1
			2	Bol_CS_hv_2
Bolgheri	Sangiovese	Pea size	1	Bol_SG_ps_1
			2	Bol_SG_ps_2
		Bunch closure	1	Bol_SG_bc_1
			2	Bol_SG_bc_2
		19 °Brix	1	Bol_SG_19_1
			2	Bol_SG_19_2
		Harvest	1	Bol_SG_hv_1
			2	Bol_SG_hv_2
	Cabernet	Pea size	1	Ric_CS_ps_1
Riccione	Sauvignon		2	Ric_CS_ps_2
		Bunch closure	1	Ric_CS_bc_1
			2	Ric_CS_bc_2
		19 °Brix	1	Ric_CS_19_1
			2	Ric_CS_19_2
		Harvest	1	Ric_CS_hv_1
			2	Ric_CS_hv_2
Riccione	Sangiovese	Pea size	1	Ric_SG_ps_1
			2	Ric_SG_ps_2
		Bunch closure	1	Ric_SG_bc_1
			2	Ric_SG_bc_2
			1	Ric_SG_19_1
		19 °Brix	I	110_00_13_1
		19 °Brix	2	Ric_SG_19_1
		19 °Brix Harvest		

where *n* base is 4,000,000.

To perform the clustering analysis, the "hits-normalizedabundance" (HNA) values were calculated as:

$$HNA = \frac{TP4M}{Hits}$$

where *TP4M* is the normalized abundance of each small RNA sequence mapping in a giving cluster and a *Hit* is defined as the number of loci at which a given sequence perfectly matches the genome.

One database was produced using the grapevine genome, and made available on the website (https://mpss.danforthcenter.org/dbs/index.php?SITE=grape_sRNA_GxE), in order to store and assist the visualization of all the sequenced libraries.

Static Clustering Analysis

The static clustering analysis was carried out as previously described by Lee et al. (2012), using a proximity-based pipeline built with custom Perl and database scripts (McCormick et al., 2011) and MySQL database queries, to group and quantify clusters of small RNAs. Briefly, the grapevine genome was divided into a series of windows of 500 bp, each window defined as a cluster. For every individual library, the small RNAs ranging from 21 to 24 nt and mapping in each cluster had their "hits-normalized-abundance" (HNA) summed up which determined the "cluster abundance." The cluster abundance was averaged for the two replicates of each library. The clusters were annotated for gene and repeat information using the V1 annotation of the reference genome (Jaillon et al., 2007; Vitulo et al., 2014), allowing the characterization of specific small RNA-producing loci.

We set a selection criterion, by which a cluster was considered as expressed when the cluster abundance was equal or greater than 30 HNA. Additionally, when investigating the ratio between two cultivars in each environment (CS/SG ratio), only those clusters where the HNA of each library in the comparison was greater than or equal to 5 (library $A \ge 5$ HNA and library $B \ge 5$ HNA) and the sum of the cluster abundance of these same libraries was higher than 30 (library A + library B > 30) were selected.

All the clustering analyses were performed using only two developmental stages for each cultivar: bunch closure was used to represent "green tissues" (g) and 19 $^{\circ}$ Brix to represent "ripened tissues" (r).

Identification of Conserved miRNAs and Prediction of Novel Candidates

The identification of annotated (conserved or known) and novel (or specie-specific) miRNAs was carried out applying a conservative and robust pipeline as described by Jeong et al. (2011) and Zhai et al. (2011), and successfully deployed in various published studies (Jeong et al., 2013; Xu et al., 2013; Arikit et al., 2014; Hu et al., 2015). Shortly, in order to recognize the conserved miRNAs, all small RNAs sequenced in the libraries were initially compared against all annotated vvi-miRNAs deposited in miRBase (version 20, Kozomara and Griffiths-Jones, 2014, http://www.mirbase.org/). Subsequently, the whole set of small RNAs passed through the five filters designed according to the properties of validated plant miRNAs and their precursors (Meyers et al., 2008), keeping track of known miRNAs throughout the filtering. The filters included, but were not limited to, minimum abundance threshold (\geq 30 TP4M), size range (18–26 nt), maximum hits to the grapevine genome (1–20), strand bias (sense/total \geq 0.9), and abundance bias [(top1+top2)/total \geq 0.7]. For each possible precursor found, the most abundant read was retained as the biologically active miRNA (also called "mature") and in cases where both the 3'-end (3p) and the 5'-end (5p) reads were highly abundant (abundance greater than 200 TP4M), the two tags were kept.

All the known vvi-miRNAs identified by the pipeline were manually inspected, to ensure that the tags identified as known miRNAs were assigned correctly to their actual precursor, and to retrieve the most abundant isoform within the tags mapping in each precursor.

Regarding the novel miRNA candidates identified using this pipeline, only those for which the most abundant tag was 20, 21, or 22 nt were retained. They were compared with all the known mature plant miRNAs in miRBase (version 20) to identify homologs. Finally, novel candidates passed through a manual evaluation of precursor secondary structures, using the plant version of the UEA sRNA hairpin folding and annotation tool (Stocks et al., 2012) and the Mfold web server (Zuker, 2003), with default settings.

miRNA Accumulation and Statistical Analysis

A miRNA was considered as "expressed" only when the values of both biological replicates were greater than or equal to the threshold set at 10 TP4M. We defined a miRNA as "vineyard-, cultivar-, or stage-specific" when it was expressed only in a given vineyard, cultivar or one specific developmental stage.

Differentially expressed miRNAs were identified using the CLCbio Genomics Workbench (v.8, Qiagen, http://www. qiagenbioinformatics.com/products/clc-genomics-workbench/) using multiple comparison analysis. We loaded the total raw redundant reads from our 48 libraries in the CLCbio package and trimmed the adaptors, considering only reads between 18 and 34 nt. We annotated miRNAs against the user defined database, comprehending our set of 122 MIRNA loci and their corresponding mature sequences. For each library, the total counts of read perfectly mapping to the miRNA precursors was considered as the input of the expression analysis.

Given the main focus of our work, we aimed at identifying miRNAs differentially expressed between the two cultivars in the same environment and developmental stage (genotypic effect), or between the three vineyards in the same cultivar and in the same developmental stage (environmental effect). For this reason, we considered each developmental stage (12 libraries) and we performed the Empirical Analysis of digital gene expression (DGE), an implementation of the "Exact Test" present in the EdgeR Bioconductor package, as implemented in CLCbio software and estimating tagwise dispersion with pairwise comparisons and setting the significance threshold to FDR-adjusted $p \leq 0.05$.

Correlation Analysis

The normalized reads (TP4M) of all miRNAs identified in this study and also the cluster abundances obtained from the static clustering analysis were submitted to another *adhoc* normalization $[\log_{10} (1+\text{TP4M}) \text{ or } \log_{10} (1+\text{HNA})]$ for correlation analysis. This normalization was chosen because of the enormous range of abundance values that produced a log-unimodal distribution and may cause significant biases in the correlation analysis when performed using TP4M or HNA values. A unity was then added to the abundance value due to the presence of zero entries. After this addition, a value of zero still corresponds to zero of the log₁₀ (1+TP4M) function, thus making consistent the comparisons among profiles.

The dendrogram was generated using the function hclust and the Pearson correlation was calculated using the function cor in R, based on the $\log_{10} (1+TP4M)$ or $\log_{10} (1+HNA)$ values for miRNAs and sRNA-generating loci respectively. Pearson's correlation coefficients were converted into distance coefficients to define the height of the dendrogram.

Heat maps were produced using MeV (MultiExperiment Viewer; Eisen et al., 1998) based on TP4M values of miRNAs abundance. The Venn diagrams were produced using the function vennDiagram in R, based on the miRNA list for each cultivar, environment and developmental stage.

Target Prediction

miRNA targets were predicted using miRferno, a built-in, plantfocused target prediction module of the software sPARTA (small RNA-PARE Target Analyzer; Kakrana et al., 2014). miRferno was run using the greedy prediction mode (tarPred H) and a seed-free system (tarScore S) for target scoring. The prediction was done in genic regions (genomeFeature 0) of the whole 12X version of the grapevine genome (Jaillon et al., 2007). The fasta file with spliced exons for each GFF transcript (V2.1.mRNA.fadownloaded from http://genomes.cribi.unipd.it/grape/) of the V2.1 annotation (Vitulo et al., 2014) was used as "feature file." To reduce the number of false positives, only targets with a prediction score value smaller than 2.5 were retained (complete range of prediction score values: 0–10).

RESULTS

High-Throughput Sequencing Statistics

Small RNA libraries were constructed and sequenced for 48 samples of grapevine berries (**Table 1**). We obtained a total of 752,020,195 raw redundant reads (Supplementary Table 1). After adaptors trimming, 415,910,891 raw clean reads were recovered, ranging from 18 to 34 nt in length (Supplementary Table 1). Eliminating the reads mapping to rRNA, tRNA, snRNA, and snoRNA sequences, 199,952,950 reads represented by 20,318,708 distinct sequences, i.e., non-redundant sequences found in the 48 libraries (Supplementary Table 1), were perfectly mapped to the *V. vinifera* PN40024 reference genome (Jaillon et al., 2007).

The libraries were analyzed to assess the size distributions of mapped reads. Distinct peaks at 21- and 24-nt (Supplementary Figure 2) were observed in all the libraries. Consistent with previous reports in grapevine (Pantaleo et al., 2010) and other

plant species (Moxon et al., 2008; Ge et al., 2013), the 21nt peak was the highest, comprising a higher proportion of redundant reads, whereas the 24-nt peak was less abundant. A few exceptions regarding the highest peak in the small RNA size profile were observed: Ric_SG_ps had the highest peak at 24nt whereas Mont_CS_ps and Mont_SG_bc did not show a clear difference between the 21- and the 24-nt peak.

Using the Pearson coefficients (Supplementary Table 2) we observed a strong association between the replicates as indicated by the high coefficients (ranging from 0.79 to 0.97).

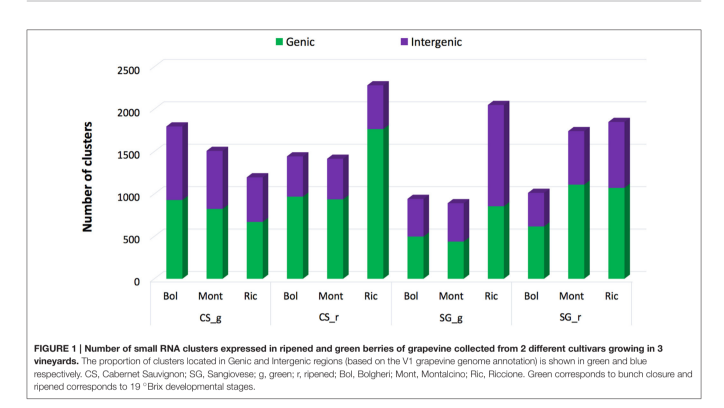
To facilitate access and utilization of these data, we have incorporated the small RNAs into a website (https://mpss. danforthcenter.org/dbs/index.php?SITE=grape_sRNA_GxE).

This website provides a summary of the library information, including samples metadata, mapped reads, and GEO accession numbers. It also includes pages for data analysis, such as quick summary of the abundances of annotated microRNAs from grapevine or other species. Small RNA-related tools are available, for example target prediction for user-specified small RNA sequences and matching criteria. Finally, and perhaps most importantly, a customized browser allows users to examine specific loci (genes or intergenic regions) for the position, abundance, length, and genomic context of matched small RNAs; with this information, coupled with the target prediction output, users can develop and assess hypotheses about whether there is evidence for small RNA-mediated regulation of grapevine loci of interest.

General Landscape of Small RNAs Distribution in Grapevine Berries in Different Environments

In order to investigate whether the overall distribution and accumulation of small RNA is affected by the interaction between different *V. vinifera* genotypes [Cabernet Sauvignon (CS) and Sangiovese (SG)] and environments [Bolgheri (Bol), Montalcino (Mont) and Riccione (Ric)], we investigated the regions in the grapevine genome from where a high number of small RNAs were being produced (sRNA-producing regions), by applying a proximity-based pipeline to group and quantify clusters of small RNAs as described by Lee et al. (2012).

The nuclear grapevine genome was divided in 972,413 adjacent, non-overlapping, fixed-size (500 bp) windows or clusters. To determine the small RNA cluster abundance, we summed the hits-normalized-abundance (HNA) values of all the small RNAs mapping to each of the 500 bp clusters, for each library (for details, see Materials and Methods). To reduce the number of false positives, we considered a cluster as expressed when the cluster abundance was greater than the threshold (HNA = 30) for a given library, eliminating regions where few small RNAs were generated, possibly by chance. Libraries from bunch closure, representing green berries, and 19 °Brix representing ripened berries, where used in this analysis. From the 972,413 clusters covering the whole grapevine genome, 4408 (0.45%) were identified as expressed (sRNA-producing regions) in at least one sample. As showed in Figure 1, CS-derived libraries have a higher number of expressed clusters when compared to SG-derived

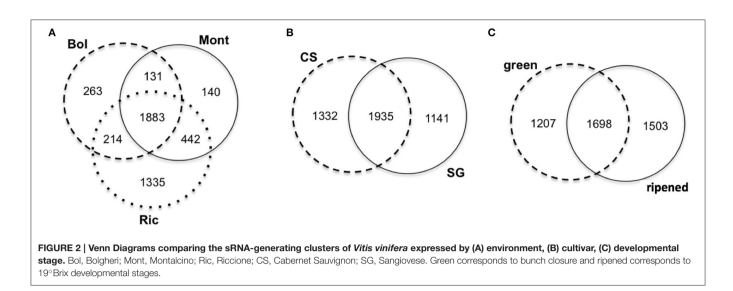


libraries of the same developmental stage and from the same vinevard. The exceptions were the Sangiovese green berries collected in Riccione and Sangiovese ripened berries collected in Montalcino, which have a higher number of expressed clusters than the respective CS ones. The two cultivars show a completely different small RNA profile across environments. When Cabernet berries were green, a higher number of sRNA-generating regions were found active in Bolgheri than in Montalcino and Riccione. Differently, ripened berries had the highest number of sRNAproducing regions expressed in Riccione, while Bolgheri and Montalcino show a similar level of expressed clusters (Figure 1). Sangiovese green berries instead show the highest number of active sRNA-generating regions in Riccione, and this number is twice the number found in Bolgheri and Montalcino that is similar. Ripened berries collected in Montalcino and Riccione show almost the same high level of sRNA-generating clusters, whereas those collected in Bogheri present a lower number (Figure 1). We also noted that when cultivated in Bolgheri, neither Cabernet Sauvignon or Sangiovese change dramatically the number of expressed clusters during ripening, while in Riccione Cabernet Sauvignon shows a 2-fold increase of sRNAproducing clusters, which is not observed in Sangiovese.

Next, the small RNA-generating clusters were characterized on the basis of the genomic regions where they map, i.e., genic, intergenic and transposable elements. In general, when the berries were green, the numbers of sRNA-generating loci located in genic and intergenic regions were roughly equal in all environments and for both cultivars, except for Sangiovese berries collected in Riccione, which show a slight intergenic disposition of sRNA-producing regions (**Figure 1**). Differently, in ripened berries on average 65% of the sRNA-generating loci were in genic regions, indicating a strong genic disposition of the sRNA-producing clusters (**Figure 1**). The shift of sRNA-producing clusters from intergenic to mostly genic is more pronounced in Cabernet Sauvignon berries collected in Riccione, with an increase of approximately 20% of expressed clusters in genic regions (**Figure 1**) when berries pass from the green to the ripened stage.

When comparing the clusters abundance among libraries, we found that 462 clusters were expressed in all libraries. The remaining 3946 expressed clusters were either shared among groups of libraries or specific to unique libraries. Interestingly, 1335 (30.3%) of the 4408 expressed clusters were specific to Riccione-derived libraries (Figure 2A). The other two environments showed a much lower percentage of specific clusters, 263 (6%) and 140 (3.2%) in Bolgheri and Montalcino respectively (Figure 2A). Comparing the expressed clusters between cultivars or developmental stages, we did not observe a similar discrepancy of specific clusters toward one cultivar or developmental stage; roughly the same proportion of specific clusters was found for each cultivar (Figure 2B) and for each developmental stage (Figure 2C). Among the 1335 specific clusters of Riccione, 605 were specific to Cabernet Sauvignon ripened berries of and 499 to Sangiovese green berries. Other smaller groups of expressed clusters were identified as specific to one cultivar, one developmental stage or also one cultivar in a specific developmental stage.

When comparing the expressed clusters with the presence of transposable elements (TE) annotated in the grapevine genome (V1), we noticed that approximately 23% of the sRNA-generating regions were TE-associated. Sangiovese green berries from Riccione have the highest proportion (26%) of TE-associated



expressed clusters, while Cabernet Sauvignon ripened berries also from Riccione show the lowest proportion (13%) of TEassociated expressed clusters. Sangiovese berries (both green and ripened) have the highest percentage of expressed clusters located in TE when cultivated in Riccione, compared to the other two vineyards. Interestingly, Cabernet Sauvignon berries show the lowest proportion of TE-associated clusters when growing in Riccione (**Figure 3A**), independently from the berry stage.

In all the libraries, Long Terminal Repeat (LTR) retrotransposons were the most represented TE. More specifically, the gypsy family was the LTR class associated with the highest number of sRNA hotspots. The other classes of TE associated with the sRNA-generating regions can be visualized in **Figure 3B**.

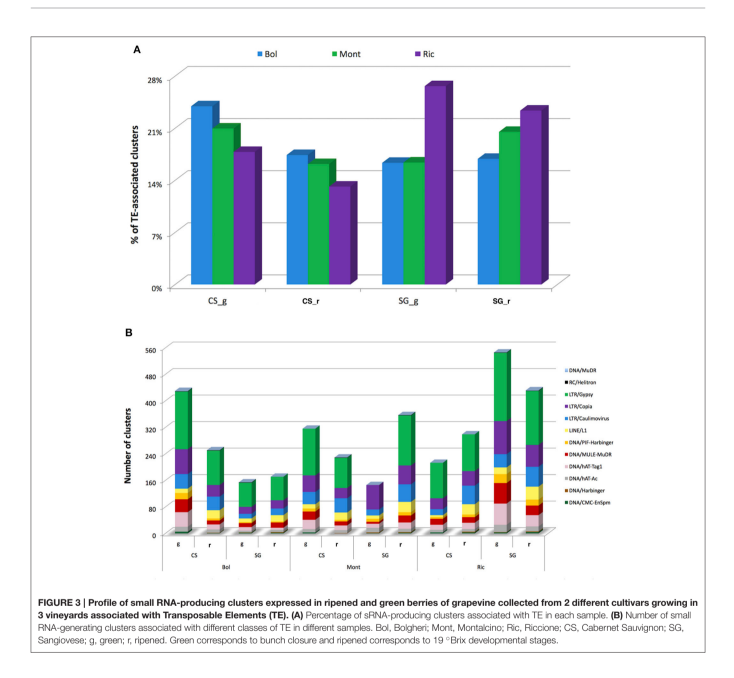
The Distribution of sRNA-Producing Loci Is Variable between the Two Cultivars, and the Level of Variation Depends on the Vineyard

To determine the global relationship of small RNA-producing loci in the different environments, cultivars and developmental stages, we performed a hierarchical clustering analysis. As showed in **Figure 4**, the libraries clearly clustered according to the developmental stage and cultivar and not according to the environments. Ripened and green berries had their profile of sRNA-generating loci clearly distinguished from each other. Inside each branch of green and ripened samples, Cabernet Sauvignon and Sangiovese were also well separated, indicating that, the cultivar and the stage of development in which the berries were sampled modulate the outline of sRNA-producing loci more than the environment.

Notwithstanding the evidence that developmental stage and variety have the strongest effect in terms of distinguishing samples clustering, we were interested to verify the environmental influence on small RNA loci expression in the two cultivars. Thus, for each sRNA-generating cluster we

calculated the ratio between cluster abundance in Cabernet Sauvignon and Sangiovese (CS/SG) in each environment and developmental stage, thereby revealing the genomic regions with regulated clusters, considering a 2-fold change threshold, a minimum abundance of 5 HNA in each library and a minimum sum of abundance of 30 HNA (library A > 5 HNA and library $B \ge 5$ HNA; library A + library B > 30 HNA; library A/library $B \ge 2$). Figure 5 shows how different environments affect the production of small RNAs. In Bolgheri, regardless the developmental stage there were many clusters with a very high abundance level in Cabernet Sauvignon (Figure 5A). In Montalcino (Figure 5B) and even more in Riccione (Figure 5C) we also observed differences between the expressions of clusters in the two cultivars, with ripened and green berries showing an almost opposite profile in terms of number of clusters more expressed in Cabernet Sauvignon or Sangiovese. When the berries were green, in Montalcino Cabernet Sauvignon shows the highest number of up-regulated clusters, while in Riccione, Sangiovese has the highest number of up-regulated clusters. The opposite behavior was noticed in ripened berries, with Sangiovese having the highest number of up-regulated clusters in Montalcino and Cabernet Sauvignon in Riccione (Figures 5B,C).

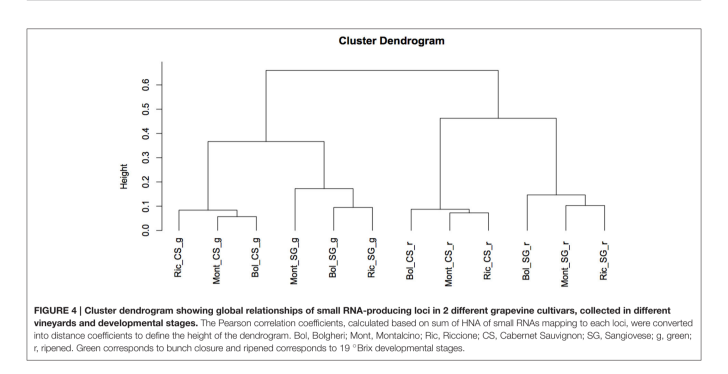
Notably, we observed a small percentage of regulated clusters (from 0.5 to 5%) exhibiting at least a 10-fold higher abundance of small RNA in Cabernet Sauvignon or Sangiovese when compared to each other (**Figures 5A–C**). An examination of those clusters showed that a substantial difference (50-fold or more) could exist between the cultivars, depending on the vineyard and the developmental stage. For example, in Riccione, a cluster matching a locus encoding a BURP domain-containing protein showed a fold change of 390 when comparing green berries of Sangiovese with Cabernet Sauvignon. The small RNAs mapping in this region were mainly 21-nt and produced from both strands (**Figure 6**). Similarly, the majority of the highly differentially expressed clusters (50-fold or more) showed a similar profile: strong bias toward 21-nt sRNAs and a low strand bias. These findings suggest that these small RNAs might be the product



of RDR (RNA-dependent RNA) polymerase activity rather than degradation products of mRNAs.

miRNAs Identification and Target Prediction

We applied a pipeline adapted from Jeong et al. (2011) and Zhai et al. (2011) to identify annotated vvi-miRNAs, their variants, novel species-specific candidates and, when possible, the complementary 3p or 5p sequences. Starting from 25,437,525 distinct sequences from all the 48 libraries, the first filter of the pipeline removed sequences matching t/r/sn/snoRNAs as well as those that did not meet the threshold of 30 TP4M in at least one library or, conversely, that mapped in more than 20 loci of the grapevine genome (considered overly repetitive to be a miRNA). Only sequences 18–26-nt in length were retained. Overall, 27,332 sequences, including 56 known vvi-miRNAs, passed through this first filter and were subsequently analyzed by a modified version of miREAP (https://sourceforge.net/projects/mireap) as described by Jeong et al. (2011). miREAP identified 1819 miRNA precursors producing 1108 unique miRNA candidates, including 47 known vvi-miRNA. Next, the sequences were submitted to the third filter to evaluate the single-strand and abundance bias retrieving only one or two most abundant miRNA sequence for each precursor previously identified. A total of 150 unique miRNA corresponding to 209 precursors were identified as candidate miRNAs. Among these 209 candidate precursors, 61 belonged to 31 known vvi-miRNA that passed all

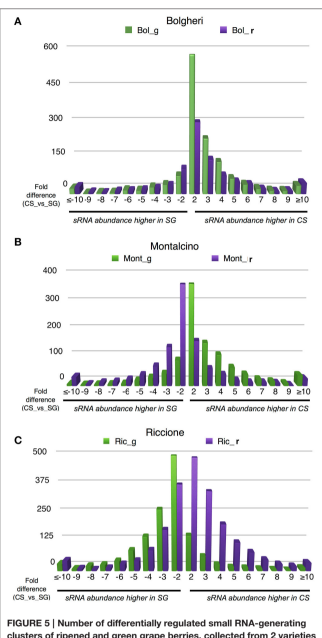


the filters and 148 were identified as putatively novel miRNA candidates. To certify that they were novel candidates rather than variants of known vvi-miRNAs we compared their sequences and coordinates with the miRNAs registered in miRBase (version 20, Kozomara and Griffiths-Jones, 2014). In order to reduce false positives and the selection of siRNA-like miRNAs, we considered only 20, 21, and 22 nt candidates whose stemloop structures were manually evaluated (see Supplementary Figure 3). Eventually, 26 miRNAs homologous to other plant species were identified with high confidence. Twenty-two were new members of nine known V. vinifera families, whereas the other four belong to three families not yet described in grapevine (Table 2). For 16 homologs we were able to retrieve also the complementary sequence. Finally, excluding these 26 miRNAs and other si-RNA like miRNAs, we identified 7 completely novel bona fide miRNAs.

Apart from the 61 known vvi-miRNAs identified by the pipeline, we searched the dataset for others known vvi-miRNAs eliminated throughout the pipeline, looking for isomiRs that were actually more abundant than the annotated sequences. Their complementary 3p or 5p sequence was also retrieved when possible. Hence 89 known vvi-miRNAs were identified in at least one of our libraries (Table 3). Among the known vvi-miRNAs identified, 24 had an isomiR more abundant than the annotated sequence and 4 have the complementary sequence as the most abundant sequence mapping to their precursor. We found 16 vvi-miRNA isomiRs that were either longer or shorter than the annotated sequence, 7 vvi-miRNAs that mapped in the precursor in a position shifted with respect to the annotated ones and one miRNA that contains a nucleotide gap when compared to the annotated sequence (Table 3). An extreme case of shifted position was found in vvi-miRNA169c, where the annotated sequence had only 5 TP4M when summing its individual abundance in the 48 libraries. Another sequence, shifted 16 bp as compared to its annotated position on the precursor had an abundance sum of 1921 TP4M, and was retained together with the annotated sequence, and named vvi-miRNA169c.1. For 36 of the 48 *V. vinifera* miRNA families deposited in miRBase we found at least one member.

An *in silico* prediction of miRNA targets was performed for the 191 mature miRNAs here identified. Using the miRferno tool (Kakrana et al., 2014), and considering only targets predicted with high stringency, 1192 targets were predicted for 143 miRNAs, including six completely novel vvi-miRNA candidates (Supplementary Table 3).

Two novel candidates (grape-m1191 and grape-m1355) seem to be involved in the regulation of important secondary metabolites biosynthesis. Among the six targets predicted for grape-m1191, the TT12 gene (transparent testa 12 -VIT_212s0028g01160) is known to be involved in the vacuolar accumulation of proanthocyanidins in grapevine (Marinova et al., 2007). For grape-m1355, 12 targets were predicted and all of them are involved in secondary metabolism pathways. Nine targets code a bifunctional dihydroflavonol 4-reductase (DFR) that is responsible for the production of anthocyanins (Davies et al., 2003), catalyzing the first step in the conversion of dihydroflavonols to anthocyanins (Boss and Davies, 2001). Another targeted gene codes a phenylacetaldehyde reductase (VIT_213s0064g00340) which, in tomato, was demonstrated to catalyze the last step in the synthesis of the aroma volatile 2-phenylethanol, important for the aroma and flavor (Tieman et al., 2007). Still this same miRNA candidate was predicted to target with high confidence (score = 0) a cinnamoyl reductaselike protein (VIT_203s0110g00350) that is part of polyphenol biosynthetic pathway (Martínez-Esteso et al., 2013). The grapem1355 candidate maps on chromosome 3, exactly on the



clusters of ripened and green grape berries, collected from 2 varieties in 3 different vineyards. The CS/SG ratios were calculated based on summed cluster abundances. (A) Data relative to berries collected in Bolgheri, (B) data relative to berries collected in Montalcino; (C) data relative to berries collected in Riccione. Bol, Bolgheri; Mont, Montalcino; Ric, Riccione; CS, Cabernet Sauvignon; SG, Sangiovese.

first exon of its target (VIT_203s0110g00350.1), in a region where another two isoforms of the same gene are located (Supplementary Figure 4). The last target of this miRNA candidate, codes a cinnamyl alcohol dehydrogenase known to be involved in the lignin biosynthesis (Trabucco et al., 2013).

Other novel vvi-miRNA candidates seem to be involved in cell proliferation (grape-m0642 targets VIT_200s0291g00090, a cyclin-related protein with hydrolase activity) and in chloroplasts-related functions (grape-m1517 targets VIT_203s0063g02020, a tic62 protein). Furthermore, for the new vvi-miRC482b candidate, besides the already known involvement of this miRNA family with disease resistance (Li et al., 2010) also predicted here, one predicted target encodes an anthocyanin 5-aromatic acyltransferase-like protein known to be involved in the biosynthesis of anthocyanin in different species (Provenzano, 2011).

As for the conserved known vvi-miRNAs, most of the well-established examples of miR-targets, such as miR156-*SPB*, miR166-*HD-ZIP*, miR171-*GRAS*, miR172-*AP2*, confirmed in several plant species and already predicted in grapevine, were also predicted here.

miRNA Accumulation among Vineyards and Genotypes

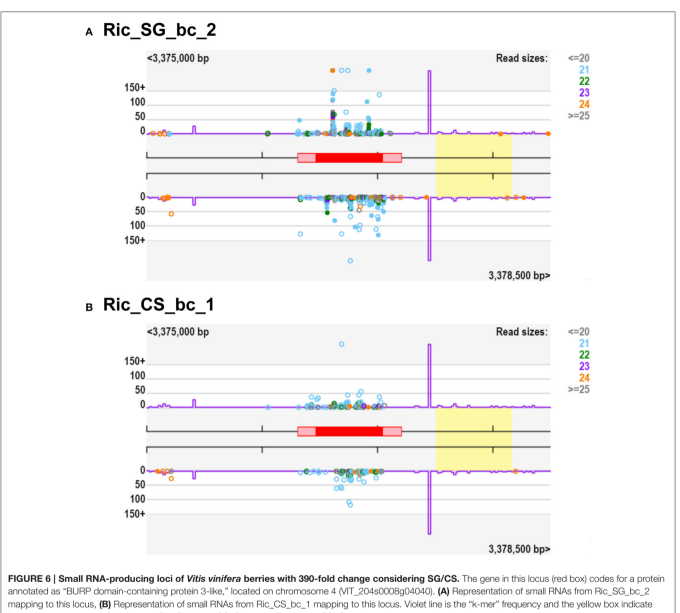
We studied miRNA profile of accumulation in the different samples. Using their normalized abundance (TP4M), i.e., their relative cloning frequency, we set an empirical cut off value equal to at least 10 TP4M in both biological replicates to consider a miRNA as expressed in a given library. Also, a miRNA was considered specific when it was expressed in one or more libraries of a unique cultivar, unique environment or unique developmental stage.

According to our established cut off, 175 miRNAs were classified as expressed in at least one of our libraries (**Figure 7**). The libraries constructed from Sangiovese berries at bunch closure collected in Bolgheri showed only 24 expressed miRNAs (**Figure 7**). For all the other libraries, expressed miRNAs ranged from 76 (Ric_SG_hv) to 148 (Ric_CS_hv) (**Figure 7**).

We found very few miRNAs specific to a given condition. The number of specific miRNAs for each cultivar, developmental stage and environment is reported in **Figures 8A–C**, respectively.

Thirty-nine vvi-miRNAs were highly expressed in almost all libraries [21 ubiquitous plus 18 expressed in all libraries except in Bol_SG_bc (**Figure 9**)], whereas other miRNAs had different accumulation patterns.

The normalized expression values of miRNAs were subjected to hierarchical clustering (HCL) and represented in a heat map (Figure 9). To examine the relatedness among cultivars, environments and developmental stages, we generated a correlation dendrogram (Figure 10). The dendrogram shows, as already suggested by the heatmaps, that a fundamental dichotomy emerges between ripened and green berries. The most evident pattern of expression is observed when comparing different developmental stages, and confirm previous observation of miRNA modulation during fruit ripening (Manning et al., 2006; Giovannoni, 2007; Carra et al., 2009; Sun et al., 2012; Cao et al., 2016). For example, some members of the vvi-miRNA156 family (f/ i and the g-5p) were highly expressed in all ripened berries, but weakly or not expressed in green berries. Differently, vvi-miR396a-3p and vvi-miR396b-3p showed the opposite profile. Similarly, vvi-miR172d, vvi-miR166b-5p, vvi-miR166f-5p, and vvi-miR396d-5p were highly expressed in green berries but weakly expressed in ripened berries and the members of the vvi-miR319 family (b/f/g and c-3p) showed a gradient of decreasing abundance from pea size to harvest.



homology to a transposable element. Ric, Riccione; CS, Cabernet Sauvignon; SG, Sangiovese; bc, green, bunch closure developmental stages.

To gain statistical evidence of miRNA differential expression driven by the environment and/or genotype, we made pairwise comparisons, keeping constant the developmental stage, and evaluating the miRNA modulation among vineyards (Montalcino vs Bolgheri vs Riccione) or between cultivars (Cabernet Sauvignon vs Sangiovese). The analyses (with an FDR ≤ 0.05) reveal that some miRNAs are differentially expressed between the two genotypes grown in the same environment, but also that a number of miRNAs are modulated by the environment. In particular the number of differentially expressed miRNAs is higher in ripened berries (19 °Brix and at harvest), while no miRNAs are differentially expressed at bunch closure stage (Supplementary Table 4). In details, 14 reads are differentially expressed at pea size stage, in at least one comparison, corresponding to 6 distinct miRNA families; 27 reads are modulated at 19 °Brix stage, corresponding to 12 miRNA families and 35 reads are differentially expressed in berries at harvest, corresponding to 12 miRNA families. It is worth noting that 4 of the 6 families modulated in the berries at pea size, are still present among the miRNAs differentially expressed in the berries sampled at 19 °Brix and at harvest (miR166, miR3627, miR477, miR3636, and miR3640), even though not always in the same comparisons.

Some of the modulated miRNAs, both novel (grape-m1355, grape-m1191) and known (miR395, miR399, and miR396) are intriguingly connected to berry development and secondary metabolism, even though most of the modulated families are still uncharacterized, or with targets not clearly involved in berry ripening and development, and deserve further studies to fully understand their biological roles.

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20 16,672,963 16,763,963 16,7240,350 17,741,733 16,141,783 17,720,353 16,3741,571 13,141,783 17,100,210 14,753 16,7270,63 13,510,210 14,753 17,533 13,510,210 14,753 14,753 14,753 14,753 14,753 14,753 119 18,87,351 18,881,468 - 7000000000000000000000000000000000000	19 12,889,963	2,890,080	+	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGCCGGTGGGGGGACC	21	83,879
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1 22,740,264 22,740,350 + TCCCTCAAAGGCTTCCATIT 21 453 19 13,141,671 13,141,788 + TCCCTCAAAGGCTTCCAATIT 21 453 19 13,510,093 13,510,210 + TCCCTCAAAGGCTTCCAATIT 21 453 19 13,510,093 13,510,210 + TCCCTCAAAGGCTTCCAATIT 21 453 19 13,510,033 13,670,210 + TCCCTCAAAGGCTTCCAATIT 21 453 19 18,872,706 18,872,824 - TCCCTCAAAGGCTTCCAATIT 21 453 19 18,81,351 18,881,468 - TCCCTCAAAGGCTTCCAATIT 21 453 11 19,755,466 19,755,588 - TCCCTCAAAGGCTTCCAATIT 21 453 11 3,865,660 3,865,686 + TCCGCTCAAAGGCTTCCAATIT 21 453 11 3,865,560 3,865,688 + TCCGCTCAAAGGCTTCCAATIT 21 453 12 3,865,560 3,865,688 +	6 19,950,730	9,950,847	I	TCCCTCAAAGGCTTCCAATTT	21	453	GCTGGAAGCCGATGGGGGGACC	21	62,642
19 13,141,671 13,141,788 + TCCCTCAAAGGCTTCCATIT 21 453 19 13,510,033 13,510,210 + TCCCTCAAAGGCTTCCAATIT 21 453 19 13,610,033 13,510,210 + TCCCTCAAAGGCTTCCAATIT 21 453 19 18,678,338 18,678,455 + TCCCTCAAAGGCTTCCAATIT 21 453 19 18,872,706 18,872,824 - TCCCTCAAAGGCTTCCAATIT 21 453 19 18,872,706 18,874,88 - TCCCTCAAAGGCTTCCAATIT 21 453 11 19,755,466 19,755,588 - TCCCTCAAAGGCTTCCAATIT 21 453 11 3,865,680 3,865,686 + TCCCTCAAAGGCTGCAAGGCTCCAATIGGGATA 20 189,848 11 3,865,560 3,865,686 + TCCGCATIGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCTGCAAGGGCGGGTGGGATA 20 189,848 10 3,865,580 3,865,686 + TCTGGCATTGGCACT 20 189,848	1 22,740,264	2,740,350	+	TCCCTCAAAGGCTTCCAATTT	21	453			
19 13,510,033 13,510,210 + TCCCTCAAAGGCTTCCAATT 21 453 19 18,678,338 18,678,455 + TCCCTCAAAGGCTTCCAATT 21 453 19 18,872,706 18,872,824 - TCCCTCAAAGGCTTCCAATT 21 453 11 19,755,466 19,755,588 - TCCCTCAAAGGCTTCCAATTGGATAGGAT 21 453 11 3,865,580 3,855,686 + TCCGCTCAAAGGCTTGCAATTGGATAGGAT 20 189,848 11 3,865,580 3,855,686 + TCCGCATTGGACCTGCACTTGGAGGAT 20 189,848 11 3,865,580 3,855,686 + TCTGGATTGGACTGCACTTGGAGGAT 21 143,513 11 3,865,580 3,855,68	19 13,141,671	3,141,788	+	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGCCGGTGGGGGGACC	21	83,879
19 18,678,338 18,678,455 + TCCCTCAAAGGCTTCCAATT 21 453 19 18,872,706 18,872,824 - TCCCTCAAAGGCTTCCAATT 21 453 19 18,873,706 18,872,824 - TCCCTCAAAGGCTTCCAATT 21 453 19 18,881,351 18,881,488 - TCCCTCAAAGGCTTCCAATT 21 453 14 19,755,466 19,755,588 - GGAATGGGCTGATTGGGATA 20 189,848 1 3,865,560 3,865,686 + CCTCGAAGGCTGCAGCT 21 453 6 17,896,112 17,896,288 + CTGGGCGGTTGCACCT 21 19,360 8 18,999,725 18,999,889 - TCTGGCATTGCACCTGCACCT 21 19,330 6 24,742,113 24,742,240 - TCTGCATTGCAGGTCACTGAGGTC 21 19,330 7 3,809,725 18,999,889 - TCTGGCATTGCACCTGCACCT 21 19,330 6 24,742,113 24,742,240 -	19 13,510,093	3,510,210	+	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGCCGGTGGGGGGACC	21	83,879
19 18,872,706 18,872,824 - TCCCTCAAAGGCTTCCAATT 21 453 19 18,81,351 18,81,468 - TCCCTCAAAGGCTTCCAATT 21 453 14 19,755,466 19,755,588 - GGAATGGGCTGATTGGGATA 20 189,848 1 3,865,560 3,865,686 + CATGGGCGGTTGGGATA 20 189,848 6 17,896,112 17,896,288 + CATGGGCGGTTGGATGGGGTAGGGGT 21 143,513 8 17,896,112 17,896,288 + TCTGGATTGGACCTGCACCT 21 19,330 8 18,999,725 18,999,889 - TCTGGATTGCACCTGCACCTGCACCT 21 19,330 8 18,999,725 18,999,889 - TCTGGATTGCACCTGCACCTGCACCT 21 19,330 6 24,742,113 24,742,240 - TTTGCACTGGTGGTCACTTAGTC 21 19,330 7 6,820,719 6,820,719 6,820,718 1,742,240 - TTACACAGGAGGAGGTCACTCAGGT 21 19,330 7<	19 18,678,338	8,678,455	+	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGTCGGTGGGGGGAAC	21	17,594
19 18,81,351 18,81,468 - TCCTCAAGGCTTCAATT 21 453 14 19,755,466 19,755,588 - GGAATGGGCTGATTGGGATA 20 189,848 1 1,9,755,466 19,755,588 - GGAATGGGCTGATTGGGATA 20 189,848 1 3,865,560 3,865,686 + CATGGGCGGTTGGGATAGGGATA 20 189,848 6 1,7996,112 17,896,288 + TCTGCATTGCACCTGCACCT 21 143,513 8 18,999,725 18,999,889 - TCTGCATTGCACCTGCACCT 21 19,330 6 24,742,113 24,742,240 - TTTGCATTGGACCTGCACCT 22 6721 5 6,820,719 6,820,833 - TTACACGGAGGAGGATGAGGGTGGTGGTGGTGGTGGTGGTGGTGGT	19 18,872,706	8,872,824	I	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGTCGGTGGGGGGACC	21	30,683
14 19.755,466 19,755,588 - GGAATGGGCTGATTGGGATA 20 189.848 1 3,865,560 3,865,686 + CATGGGCGGTTTGGTAAGAGG 21 143,513 6 17,896,112 17,896,288 + CATGGGCGGTTTGGTAAGAGG 21 19,330 7 8 18,999,725 18,999,889 - TCTGCATTTGCACCTGCACCT 21 19,330 5 24,742,113 24,742,240 - TTTGCAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	19 18,881,351	8,881,468	I	TCCCTCAAAGGCTTCCAATTT	21	453	GTTGGAAGCCGGTGGGGGGACC	21	83,879
1 3,865,560 3,865,686 + CATGGGCGGTTTGGTAAGAGG 21 143,513 6 17,896,112 17,896,288 + TCTGCATTTGCACCTGCACCT 21 19,330 8 18,999,725 18,999,889 - TCTGCATTGCACCTGCACCT 21 19,330 6 24,742,113 24,742,240 - TTTGCATTGCACCTGCACCT 21 19,330 7 5 6,820,719 6,820,833 - TTTGCAGGGGGGTGGGTCACGGGGG 22 6721 17 3,771,923 3,772,046 + GAGGTGATCAGCAAGAG 21 31 29 1,474,782 1,474,863 + GAGGTGATCAGCAAGAG 21 31	14 19,755,466	9,755,588	I	GGAATGGGCTGATTGGGATA	20	189,848	TTCCCAATGCCGCCCATTCCAA	22	17,229
6 17,896,112 17,896,288 + TCTGCATTTGCACCTGCACCT 21 19,330 8 18,999,725 18,999,889 - TCTGCATTTGCACCTGCACCT 21 19,330 5 24,742,113 24,742,240 - TTTTGTGCTGGGGGGGGGGGGG 22 6721 6 6 6,820,719 6,820,833 - TTTGCAGGGGGGGGGGGG 22 6721 17 3,771,923 3,772,046 + GAGGTGATCAGCAAGGAG 21 31 29 1,474,782 1,474,863 + GAGGTGATCAGCAAGGAG 21 31	1 3,865,560	3,865,686	+	CATGGGCGGTTTGGTAAGAGG	21	143,513	TCTTACCAACACCTCCCATTCC	22	37,745
8 18,999,725 18,999,889 - TCTGCATTGCACCTGCACCT 21 19,330 5 24,742,113 24,742,240 - TTTGTGCTGGTCGTCGTCTGGTC 22 6721 6 6,820,719 6,820,719 6,820,833 - TTACACAGAGAGATGACGGTGG 22 6721 17 3,771,923 3,772,046 + GAGGTGATCAGCAAGAG 21 31 29 1,474,782 1,474,863 + GAGGTGATCAGCAAGAG 21 31	6 17,896,112	7,896,288	+	TCTGCATTTGCACCTGCACCT	21	19,330	AGGTGCAGGTGAAGGTGCAGA	21	207
5 24,742,113 24,742,240 - TTTGTGGGTGGTCAGTC7AGTC 22 6721 5 6,820,719 6,820,833 - TTACACAGAGAGATGACGGTGG 22 535 1 17 3,771,923 3,772,046 + GAGGTGATCAGCAGAGA 21 31 29 1,474,782 1,474,863 + GAGGTGATCAGCAAGAG 21 31	8 18,999,725	8,999,889	I	TCTGCATTTGCACCTGCACCT	21	19,330			
5 6,820,719 6,820,833 - TTACACAGAGAGATGACGGTGG 22 535 . 17 3,771,923 3,772,046 + GAGGTGATGATCAGCAAGAG 21 31 29 1,474,782 1,474,863 + 4	5 24,742,113	4,742,240	I	TITTGTTGCTGGTCATCTAGTC	22	6721	TTAGATGATCATCAACAAACA	21	12,436
17 3,771,923 3,772,046 + GAGGTGATAGATCAGCAAGAG 21 31 29 1,474,782 1,474,863 +	5 6,820,719	3,820,833	I	TTACACAGAGAGATGACGGTGG	22	535	ACCGTCTTTCTCTGTATAAGC	21	1227
29 1,474,782 1,474,863 +	17 3,771,923	3,772,046	+	GAGGTGATAGATCAGCAAGAG	21	31			
	29 1,474,782	1,474,863	+				CCCGAGAGGACTTAGTGGATC	21	413
grape-m1191 29 469,980 470,068 - TTGCTGAACA	29	470,068	ī				TTGCTGAACAAGAGAGAACCT	21	1859
grape-m1355 3 13,241,508 13,241,588 – GCCGCGTTGGAGCAGGAGCTT 21 740	ю Т	3,241,588	I	GCCGCGTTGGAGCAGGAGCTT	21	740			
grape-m1517 5 11,023,593 11,023,756 – AACTACTGAAT	5 11,023,593	1,023,756	I				AACTACTGAATCATTGACCAG	21	06
6,902,214 6,902,359 –	6 6,902,214	3,902,359	I				GGCACGACAGTCTTGGACGCC	21	31
grape-m1738 8 21,905,716 21,905,814 – TCGTAGTGGCTGCGACAGCTCC 22 2406	8 21,905,716	1,905,814	ī	TCGTAGTGGCTGCGACAGCTCC	22	2406			

Family	miRNA	Chr ^a	Start ^a	End ^a	Strand ^a	5p Sequence	nt ^b	Abundance 5p ^c	3p Sequence	nt ^b	Abundance 3p ^c
wi-miR156	wi-miR156b	4	5,357,061	5,357,310	I				TGACAGAAGAGAGTGAGCAC	20	10,561
	wi-miR156c	4	848,207	848,307	I	TGACAGAAGAGAGTGAGCAC	20	10,561			
	wi-miR156d	11	7,623,202	7,623,334	I	TGACAGAAGAGAGTGAGCAC	20	10,561	TGCTCACCTCTCTTTCTGTCAGC	23	1089
	wi-miR156e	11	1,504,195	1,504,301	I	TGACAGAGGAGAGTGAGCAC	20	234			
	wi-miR156f	14	26,463,671	26,463,773	+	TTGACAGAAGATAGAGGAGCAC	21	26,445			
	wi-miR156g	17	3,046,310	3,046,441	I	TTGACAGAAGATAGAGAGCAC	21	26,445	GCTCTCTAGACTTCTGTCATC	21	901
	wi-miR156i	14	19,727,139	19,727,358	I				TTGACAGAAGATAGAGAGCAC	21	26,445
vvi-miR159	wi-miR159c	17	2,609,190	2,609,409	Т	GAGCTCCTTGAAGTCCAATAG	21	1644	TITGGATTGAAGGGAGCTCTA	21	198,072
wi-miR162	wi-miR162	17	4,716,504	4,716,636	+	GGATGCAGCGGTTCATCGATC	21	365	TCGATAAACCTCTGCATCCAG	21	130,641
vvi-miR164	wi-miR164a	7	3,287,472	3,287,590	I	TGGAGAGCAGGGCACGTGCA	21	198			
	wi-miR164c	œ	1,008,0445	10,080,636	+	TGGAGAGCAGGGCACGTGCA	21	198	CATGTGCCCCTCTTCCCCCATC	21	976
	wi-miR164d	14	1,414,567	1,414,682	I	TGGAGAGCAGGGCACGTGCA	21	198			
wi-miR166	wi-miR166a	00	3,302,784	3,302,939	I	AATGAGGTTTGATCCAAGATC	21	1886	TCTCGGACCAGGCTTCATTCC	21	1,172,737
	wi-miR166b	12	17,937,384	17,937,510	+	TCGGACCAGGCTTCATTCCTC	21	10,941	GGAATGTTGGCTGGCTCGAGG	21	120,762
	wi-miR166c	15	16,978,558	16,978,745	I	GGAATGTTGTCTGGCTCGAGG	21	54,367	TCGGACCAGGCTTCATTCCCC	21	17,706,997
	wi-miR166d	16	21,405,202	21,405,388	I	GATTGTTGTCTGGCTCGAGGC	21	2661	TCGGACCAGGCTTCATTCCCC	21	17,706,997
	wi-miR166e	0	2,255,708	2,255,901	+	GGAATGTTGTCTGGCTCGAGG	21	54,367	TCGGACCAGGCTTCATTCCCC	21	17,706,997
	wi-miR166f	7	19,450,000	19,450,127	+	GGAATGTTGGCTGGCTCGAGG	21	10,941	TCGGACCAGGCTTCATTCCCC	21	17,706,997
	wi-miR166g	7	453,844	453,971	I	GGAATGTTGTCTGGTTCGAGA	21	262	TCGGACCAGGCTTCATTCCCC	21	17,706,997
	wi-miR166h	£	6,741,189	6,741,288	I				TCGGACCAGGCTTCATTCCCC	21	17,706,997
vvi-miR167	wi-miR167b	14	7,137,373	7,137,501	+	TGAAGCTGCCAGCATGATCT-	21	1143	AGATCATGTGGCAGTTTCACC	21	536
	wi-miR167d	20	7,490,493	7,490,606	+	TGAAGCTGCCAGCATGATCT-	21	1143			
	wi-miR167e	Ð	5,845,370	5,845,489	+	TGAAGCTGCCAGCATGATCT-	21	1143	AGATCATGTGGCAGTTTCACC	21	536
wi-miR168	wi-miR168	N	17,944,786	17,944,947	I	TCGCTTGGTGCAGGTCGGGAA	21	43,158	CCCGCCTTGCATCAACTGAAT	21	3558
	wi-miR169c	4	2,265,913	2,266,028	I	CAGCCAAGGATGACTTGCCGG	21	5			
vvi-miR169	wi-miR169c.1	4	2,265,913	2,266,028	I	TGTAGGGAGTAGAATGCAGC	20	1921			
	wi-miR169e	14	25,082,574	25,082,756	I	TAGCCAAGGATGACTTGCCT-	22	3136			
	wi-miR169g	Ø	21,104,448	21,104,568	+	CAGCCAAGGATGACTTGCCGA	21	13	*CCGGCAAGTTGTCTTTGGCTAC	23	151
	wi-miR169r	11	16,415,128	16,415,239	+	TGAGTCAAGGATGACTTGCCG	21	11	*GGCAAGTTGACTTGACTCAGT	22	1181
	wi-miR169t	11	16,399,564	16,399,676	+	CGAGTCAAGGATGACTTGCCGA	22	14	*GGCAAGTTGACTTGACTCAGT	22	1181
wi-miR171	wi-miR171b	12	5,542,396	5,542,529	I				-TTGAGCCGCGTCAATATCTCC	24	1068
	wi-miR171i	17	893,536	893,632	+				TGATTGAGCCGTGCCAATATC	21	811
vvi-miR172	wi-miR172d	œ	12,667,173	12,667,279	+				TGAGAATCTTGATGATGCTGC-	23	6451
wi-miR2111	wi-miR2111	Ø	15,368,664	15,368,748	I	TAATCTGCATCCTGAGGTCTA	21	1476	GTCCTCTGGTTGCAGATTACT	21	95
vvi-miR2950	wi-miR2950	7	14,340,406	14,340,517	+	TTCCATCTTGCACACTGGA	21	3176	TGGTGTGCACGGGGATGGAATA	21	581
wi-miR319	wi-miR319b	-	4,189,556	4,189,753	+				TTGGACTGAAGGGGAGCTCCC-	21	9157
		¢						T C T			

(Continued)

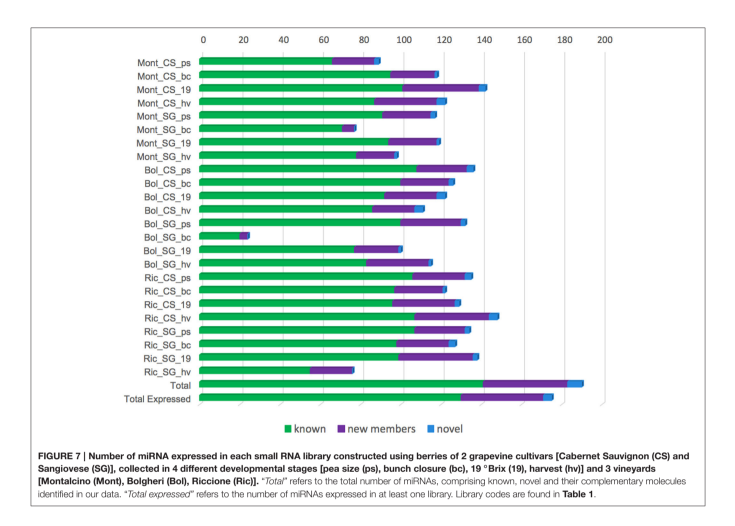
Family	miRNA	Chr ^a	Start ^a	End ^a	Strand ^a	5p Sequence	nt ^b	Abundance 5p ^c	3p Sequence	nt ^b	Abundance 3p ^c
	wi-miR319e	11	4,317,223	4,317,329	+				TTTGGACTGAAGGGAGCTCCT	21	40,107
	wi-miR319f	9	9,137,252	9,137,445	+				TTGGACTGAAGGGAGCTCCC-	21	9157
	wi-miR319g	17	3,675,979	3,676,209	I				TTGGACTGAAGGGAGCTCCC-	21	9157
wi-miR3623	wi-miR3623	18	24,650,004	24,650,151	+	TCACAAGTTCATCCAAGCACCA	22	108,278	GTGCTTGGACGAATTTGCTAAG	22	20,404
wi-miR3627	wi-miR3627	14	28,302,559	28,302,681	I	TTGTCGCAGGAGAGACGGCACT	Γ 22	2444	TCGCCGCTCTCCTGTGACAAG	21	12,342
vvi-miR3629	wi-miR3629a	13	18,253,987	18,254,170	+	CGCATTITCTCAGCAGCCAAG	21	÷	TGGCTGCTGAGAAAATGTAGG-	22	371
	wi-miR3629c	17	822,209	822,338	+	TGGCTGCTGAGAAAATGTAGG-	22	371			
wi-miR3632	wi-miR3632	14	23,394,889	23,395,015	+	GGATTGGGGGGCCGATGGAAAGG	3 22	887	TTTCCCAGACCCCCCAATACCAA	22	1578
vvi-miR3633	wi-miR3633a	17	5,521,913	5,522,060	+	GGAATGGATGGTTAGGAGAG	20	45,786	TTCCTATACCACCCATTCCCTA	22	1469
	wi-miR3633b	17	5,521,557	5,521,691	+	GGAATGGGTGGCTGGGATCT	20	10,270	GTTCCCATGCCATCCATTCCTA	22	5610
wi-miR3634	wi-miR3634	17	5,681,202	5,681,314	T	GGCATATGTGTGACGGAAAGA	21	1563	TITCCGACTCGCACTCATGCCGT	23	753,254
vvi-miR3635	wi-miR3635	18	27,357,619	27,357,780	+	GGCATGTGTGGGGGCATAATAG	21	289	ATTATGTCCCACACATGCCTC	21	399
wi-miR3636	wi-miR3636	16	5,414,817	5,414,967	I	TCGGTTTGCTTCTTTGATAGATTC	0 24	352	1	24	4495
									TGTCGGAGAAGCAAGTCGGAG <mark>AGT</mark>		
vvi-miR3640	wi-miR3640	16	11,986,842	11,987,014	+	ACCTGATTGGTGATGCTTTTTGG 24	G 24	1321	 AAAGGCATCATCAATCAGGTAATG	24	1203
wi-miR390	wi-miR390	9	8,159,519	8,159,657	+	AAGCTCAGGAGGGGATAGCGCC	21	5046			
vvi-miR393	wi-miR393a	16	17,247,172	17,247,327	T	TCCAAAGGGATCGCATTGATC	21	5152	ATCATGCTATCCCTTAGGAAC	21	775
	wi-miR393b	13	4,265,132	4,265,213	+	TCCAAAGGGATCGCATTGATC	21	5152			
wi-miR394	wi-miR394a	12	17,122,005	17,122,092	I	TTGGCATTCTGTCCACCTCC-	22	412			
	wi-miR394b	18	1,413,038	141,3130	I	TTGGCATTCTGTCCACCTCC	20	412			
vvi-miR395	wi-miR395a	÷	6,527,921	6,528,019	+				CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395c	-	6,499,899	6,500,020	I	GTTCCCTTGACCACTTCACTG	21	5850	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395d	-	6,512,760	6,512,848	+				CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395e	-	6,505,233	6,505,354	+	GTTCCCTTGACCACTTCACTG	21	5850	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395f	-	6,489,527	6,489,642	+	GTTCCCCTGACCACTTCACTG	21	4729	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395h	-	6,566,637	6,566,758	+	GTTCCCTTGACCACTTCACTG	21	5850	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395i	-	6,562,627	6,562,742	+	GTTCCCCTGACCACTTCACTG	21	4729	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR395j	-	6,553,011	6,553,126	+	GTTCCCCTGACCACTTCACTG	21	4729	CTGAAGTGTTTGGGGGGAACTC	21	6839
	wi-miR395k	-	6,536,764	6,536,885	+	GTTCCCTTGACCACTTCACTG	21	5850	CTGAAGTGTTTGGGGGGAACTC	21	6899
	wi-miR3951	-	6,559,083	6,559,199	+	GTTCCCCTGACCACTTCACTG	21	4729	CTGAAGTGTTTGGGGGGAACTC	21	6899
wi-miR396	wi-miR396a	6	7,372,499	7,372,649	I	TTCCACAGCTTTCTTGAACTA	21	15,069	AAGAAAGCTGTGGGGGGGGGACATGGC	C 24	2147
	wi-miR396b	11	5,246,778	5,246,913	+	TTCCACAGCTTTCTTGAACTT	21	85,683	GTTCAAGAAAGCTGTGGGGAAA	21	6917
	wi-miR396c	4	5,119,592	5,119,698	I	TTCCACAGCTTTCTTGAACTG	21	3900			
	wi-miR396d	11	5,253,095	5,253,244	I	TTCCACAGCTTTCTTGAACT-	21	7387	GTTCAATAAGCTGTGGGGAAG	21	1899
vvi-miR397	wi-miR397a	20	11,971,890	11,972,015	Ι	TCATTGAGTGCAGCGTTGATG	21	432			
wi-miR398	wi-miR398b	9	16,503,544	16,503,633	I				TGTGTTCTCAGGTCGCCCCTG	21	16,121
	wi-miR398c	9	15.575.579	15.575.668	+				TGTGTTCTCAGGTCGCCCCTG	21	16.121

Family	miRNA	Chr	Start ^a	End ^a	Strand ^a	pb Sequence		Abundance 5p ^c	3p Sequence	ut ²	
vvi-miR399	wi-miR399a	10	2,989,435	2,989,561	+	GTGTGAITCTCCTTTGGCAGA	21	503	TGCCAAAGGAGAATTGCCCTG	21	1289
	wi-miR399b	16	15,618,708	15,618,845	I				TGCCAAAGGAGAGAGTTGCCCTG	21	2996
	wi-miR399c	15	15,232,197	15,232,281	+				TGCCAAAGGAGAGATTGCCCTG	21	2996
	wi-miR399e	10	2,992,220	2,992,341	Ι				TGCCAAAGGAGATTTGCCCGG	21	184
	wi-miR399h	10	2,983,543	2,983,634	+				TGCCAAAGGAGAATTGCCCTG	21	1289
	wi-miR399i	0	4,101,786	4,101,937	+	GGGCTTCTCCCTTCTGGCAGG	22	148	CGCCAAAGGAGAGTTGCCCTG	21	113,055
wi-miR403	wi-miR403a	Ð	65,247	65,361	+				TTAGATTCACGCACAAACTCG	21	82,567
	wi-miR403b	Ð	600,176	600,266	+				TTAGATTCACGCACAAACTCG	21	82,567
	wi-miR403d	ß	166,467	166,582	+	AGTITGTGCGCGAATCCAACC	21	201	TTAGATTCACGCACAAACTCG	21	82,567
	wi-miR403e	5	168,096	168,213	+				TTAGATTCACGCACAAACTCG	21	82,567
	wi-miR403f	7	4,179,658	4,179,795	I	AGTITGTGCGTGACTCTAAAA	21	523	TTAGATTCACGCACAAACTCG	21	82,567
vvi-miR408	wi-miR408	7	5,011,920	5,012,056	+	CGGGGACGAGGTAGTGCATGG	21	3223	ATGCACTGCCTCTTCCCTGGC	21	6566
wi-miR477	wi-miR477b	2	1,237,529	1,237,808	+	ACTCTCCCTCAAGGGCTTCT-	22	513	CGAAGTCTTTGGGGGGGGAGAGTGG	21	3289
						Ű					
vvi-miR479	wi-miR479	16	21,573,744 21,573,866	21,573,866	+	TGTGGTATTGGTTCGGCTCATC	22	4449	CGAGCCGAACCAATATCACTC	21	46,073
wi-miR482	wi-miR482	17	5,523,009	5,523,155	+	*AATTGGAGAGTAGGAAAGCTT	22	139,166	TCTTTCCTACTCCTCCCATTCC	22	42,801
vvi-miR535	wi-miR535a	25	1,392,255	1,392,353	I	TGACAACGAGAGAGAGCACGC	21	18,893			
	wi-miR535c	25	1,346,353	1,346,483	I	TGACAACGAGAGAGAGCACGC	21	18,893	GTGCTCTCTGTCGCTGTCATA	21	2191

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TABLE 3 | Continued

—Missing nucleotide. Nucleotides that differ from the annotated sequence are shown in red. *Indicates that the most abundant sequence in our dataset does not correspond to the mature annotated miRNA, but to the star sequence.



DISCUSSION

Using high throughput sequencing coupled with robust bioinformatics pipelines we analyzed small RNAs derived from the berries of Cabernet Sauvignon and Sangiovese, grown sideby-side in three vineyards, representative of different grapevine cultivation areas in Italy (Bolgheri, Montalcino, and Riccione). We obtained nearly 750 MB reads comprising a significant proportion of small (21–24 nt) RNAs. The size distribution profiles of our libraries were in general consistent with previous reports in berry grapevine, where the 21-nt class was more abundant than the 24-nt class (Pantaleo et al., 2010; Wang et al., 2012; Han et al., 2014; Kullan et al., 2015).

Our analysis revealed dynamic features of the regulatory network mediated by miRNAs and other small RNAs, at the basis of genotype-environment interactions.

Genotype and Environment Effects on Small RNA Profiles

Plants evolved a series of pathways that generate small RNAs of different sizes with dedicated functions (Vazquez, 2006; Khraiwesh et al., 2012). Although the various small RNA classes have been intensively studied, we are still far from understanding

how many small RNA pathways exist, and how they are connected (Vazquez, 2006). Additionally, new classes of small non-coding RNAs continue to be discovered and many studies demonstrate a substantial redundancy and cross-talk between known small RNA pathways (Agarwal and Chen, 2009; Ghildiyal and Zamore, 2009; Bond and Baulcombe, 2014; Harding et al., 2014). Estimating the exact percentage of the plant genome covered by small RNA-generating loci still remains a challenge.

By applying static cluster analysis, we investigated small RNA abundances across the genome, identifying 4408 small RNAs producing hotspots. We analyzed their expression in different cultivars, environments and developmental stages, highlighting that the majority of the considered small RNA producing regions was modulated in different conditions. This suggests a strong influence of small RNAs in the response to environment in grapevine berries. Only 462 small RNA-generating loci, corresponding to about 10% of the total, were expressed in all the analyzed libraries, possibly involved in essential biological pathways.

Comparing the two cultivars, we observed, with few exceptions, that Cabernet Sauvignon berries have a higher number of expressed sRNA-generating loci than Sangiovese berries (**Figure 1**) when collected in the same conditions (i.e.,

vineyard and developmental stage). Considering the fact that small RNAs are implicated in the regulation of gene expression in several processes (Chen, 2009; Trindade et al., 2011), the higher number of small RNAs expressed in Cabernet Sauvignon compared to Sangiovese berries may reflect a buffering effect of small RNAs influencing grapevine response to diverse growing environments. We believe that these characteristics may have contributed to the wide diffusion of Cabernet Sauvignon, allowing its wide cultivation in almost all wine producing countries. This is not the case for Sangiovese whose cultivation is more restricted. It is worth noting that Sangiovese is considered a very unsettled grapevine cultivar (Poni, 2000), showing a wide range of variability in response to year, clone and bunch exposure (Rustioni et al., 2013). Differently, Cabernet Sauvignon is a cultivars showing less inter-annual differences in terms, for example, of concentration of secondary metabolites (Ortega-Regules et al., 2006).

SG

To better evaluate varietal differences in response to the environment, we calculated the CS/SG ratio for the small RNA producing hotspots in the three vineyards. An interesting example is found in green berries sampled in Riccione. A region on chromosome 4 (3,376,501-3,377,000) showed a 390-fold change in the small RNA abundance, when comparing Cabernet vs. Sangiovese (Figure 6). Most of the reads produced in this region are 21 nt long and are also phased in intervals of 21 nt from both strands, typical of a phased locus (PHAS). The gene in this locus, also known as VvRD22g, encodes a BURP domaincontaining protein, involved in an ABA-mediated abiotic stress response, which persists still after long periods of stress (Matus et al., 2014). The small RNAs profile suggests that the locus is regulated by *phased* siRNAs similarly to the mechanisms already described for PPR, NB-LRR, and MYB gene families (Howell et al., 2007; Zhai et al., 2011; Xia et al., 2013; Zhu et al., 2013). This is a clear example of GxE interactions since the BURP domain gene modulates *phased* siRNAs production in the two cultivars only when grown in Riccione.

When removing the threshold of minimum cluster abundance set to 5 HNA, in the CS/SG ratio, a high number of clusters

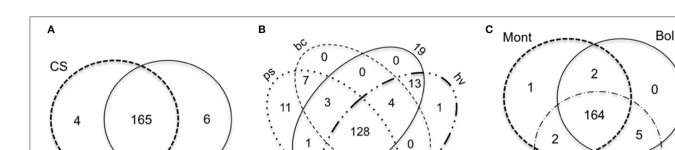
(ranging from 70 to 370 depending on the sample analyzed) with fold change greater than 50 was found, where one of the libraries has 0 HNA and the other any number greater than 30 HNA. This fact suggests a very strong modulation of the expression of small RNAs between the two cultivars, which is more or less pronounced depending on the vineyard where the berries were cultivated. A similar situation was observed comparing the expression level of small RNAs between reciprocal hybrids of *Solanum lycopersicum* and *S. pimpinellifolium* (Li et al., 2014).

The ripening process of grapevine berries is highly affected by the environment (van Leeuwen et al., 2004, 2007) and we observed the impact of the environment on the ripening process in the expression of small RNAs. The most relevant observation is that Riccione is very peculiar in relation to the activation of sRNA hotspots, as indicated by the high number of Riccionespecific clusters (Figure 2A) and by the extreme modification it induces in the CS/SG ratio (Figure 5): in Riccione in fact this ratio decreases in green berries and increases in ripened berries, and this is not observed in any other vineyard; in addition to this the already discussed example of BURP domain gene, is observed in Riccione, as well. Riccione is the most diverse environment when compared to Montalcino and Bolgheri. Riccione is located at the Adriatic coast and has a temperate sub-littoral climate, while Montalcino and Bolgheri are both located in Tuscany with typically Mediterranean climate.

Moreover, both cultivars show a peculiar profile of small RNA loci during berries ripening, in Riccione. The expression of small RNA loci in Cabernet Sauvignon berries drastically changed during development, especially when collected in Riccione (**Figure 1**), not only in the number of active loci but also in the different genic or intergenic disposition: ripened berries have a 2.6-fold increase in small RNA loci active in genic regions. Differently, when Sangiovese is grown in Riccione, there is a very high number of small RNA loci active in green berries, mainly associated to transposable elements that remains almost stable during development although the proportion of intergenic loci is reduced. Sangiovese berries collected in Montalcino show a 2.5fold increase of small RNA producing loci during development.

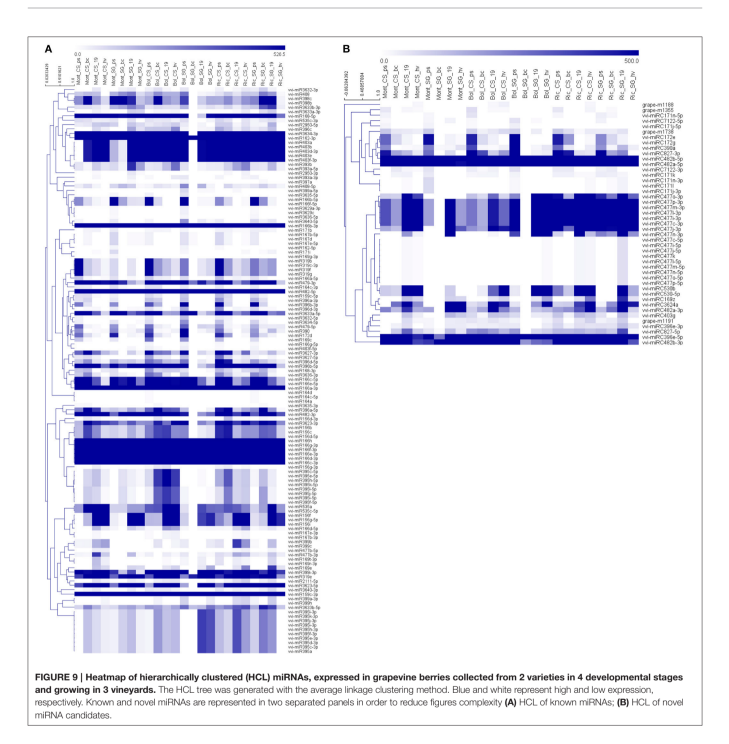
1

Ric



0

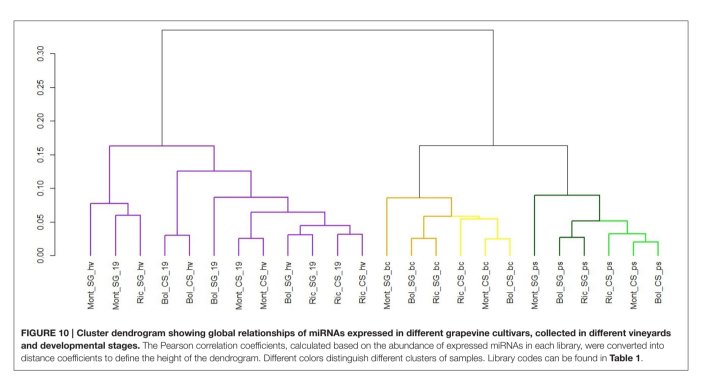
FIGURE 8 | Venn diagrams comparing the miRNAs expressed in small RNA libraries of Vitis vinifera constructed using berries. The venn diagrams are divided by (A), cultivar; (B), developmental stage; (C), environment. Bol, Bolgheri; Mont, Montalcino; Ric, Riccione; CS, Cabernet Sauvignon; SG, Sangiovese.



Differences during berry development between the cultivars may explain their different behavior in different environments, and the characteristics of each vineyard may favor one or other variety according to their demands. For example, Sangiovese needs a long growing season (it is slow to ripen) with sufficient warmth to fully ripen (Poni, 2000). Consequently, cooler environments will require a reprograming of Sangiovese gene expression in order to achieve ripening. Other factors such as composition of soil, level of humidity, photoperiod and density of cultivation may be exerting the same influence on the ripening of the berries triggering the activation of different small RNA loci.

miRNAs Expression Is Mainly Dependent on the Developmental Stage but a Few miRNAs Are Directly Modulated by the Vineyard and the Cultivar

Applying a conservative pipeline to the analysis of our 48 small RNA libraries, we recognized 89 known and annotated grapevine



miRNAs. In addition, when compared to previous reports in grapevine (Alabi et al., 2012; Han et al., 2014; Wang et al., 2014) we identified 7 completely novel miRNAs plus 26 homologous to other plant species, but novel to grapevine. This is a remarkable number considering the stringency of our pipeline and that our study is based only on four developmental stages of berries.

The outline of miRNA accumulation across samples is different from that of sRNA-producing loci. While the expression of sRNA-generating regions allows distinguishing very well between ripened and green berries and also between cultivars (**Figure 4**), the accumulation of miRNAs shows a clear distinction only between ripened and green berries, and when the berries were green, we observe a further dichotomy separating the two cultivars and the two green developmental stages. The same pattern of miRNA accumulation among green and ripened berries of grapevine (cv. Corvina) was observed when we described the miRNA expression atlas of *Vitis vinifera* (Kullan et al., 2015).

Comparing the distribution of miRNAs expressed throughout our samples, we found a set of 39 miRNAs ubiquitous (21) or nearly ubiquitous (18) to all the libraries, and very few miRNAs specific of a cultivar, vineyard or developmental stage. All these 39 miRNAs belong to known vvi-miRNA families. With few exceptions, the same set of miRNAs was also found expressed in all the small RNA libraries constructed with different tissues of the grapevine cv. Corvina (Kullan et al., 2015), where the population of expressed miRNAs appears highly variable apart from a well-defined group of miRNAs, probably related to the basal metabolism. These findings are also consistent with previous report in grapevine where a small number of known tissue-specific miRNAs was described (Wang et al., 2014).

Considering the ripening process as shown in the heat maps (**Figure 9**), and the correlation dendrogram, it is clear that most miRNAs are modulated during the developmental process.

For some miRNA families, we observed the same peculiar patterns of miRNA accumulation, previously described in the grapevine miRNA atlas (Kullan et al., 2015), e.g., an increase of accumulation toward ripening for miR156 f/g/i, and a decrease for miR166c/e, miR172d, miR319, and miR396a/b, but this is not the main focus of our paper.

To establish genotype and environmental influence on miRNA modulation, we performed a statistical analysis that revealed a number of miRNAs differentially expressed. Being aware of the fact that we had only two biological replicates, we applied the exact test as implemented in the EdgeR package. This test has been recently judged a very robust tool that can be used in experiments similar to our, because of its low false positive rate and relative high true positive rate in the presence of a fold change higher than 4 (Schurch et al., 2016).

Considering berries at the same developmental stages, we compared Sangiovese vs. Cabernet Sauvignon in a given vineyard and Montalcino vs. Bolgheri, Montalcino vs. Riccione, and Bolgheri vs. Riccione keeping the cultivar fixed. In total we performed 9 pairwise comparisons for each developmental stage. In general, we observed that berries at 19 °Brix and at harvest show a higher number of differentially expressed miRNAs.

The most interesting examples are represented by two novel miRNAs, whose predicted targets are related to the biosynthesis and accumulation of secondary metabolites, which are of crucial importance in grapevine berries, since its quality depends mainly on its metabolites (Ali et al., 2010). The candidate grape-m1191 is differentially expressed in Sangiovese between Riccione and Bolgheri (Ric_SG_19 vs. Bol_SG_19) and was predicted to target the transparent-testa 12 gene (VIT_212s0028g01160) that encodes a multidrug secondary transporter-like protein (MATE) involved in the vacuolar accumulation of the flavonoid proanthocyanidin in different species including grapevine (Debeaujon et al., 2001; Bogs et al., 2007; Marinova et al.,

2007; Zhao et al., 2010). Also, in grapevine some studies provide evidences that the intracellular transport of acylated anthocyanins is catalyzed by a MATE transporter (Gomez et al., 2009; He et al., 2010).

The grape-m1355 seems to be involved in four different pathways, all related to secondary metabolites. It is differentially expressed in Montalcino between the two varieties (Mon_CS_hv vs. Mon_SG_hv) and was predicted to target a cinnamoyl reductase-like protein (CCR) (VIT_203s0110g00350), which is part of the of the polyphenol biosynthetic pathway (Leple et al., 2007); a cinnamyl alcohol dehvdrogenase (VIT_206s0004g02380) involved in the lignin biosynthesis (Trabucco et al., 2013); a phenylacetaldehyde reductase (VIT_213s0064g00340), which catalyzes, in tomato, the last step in the synthesis of the volatile 2-phenylethanol, important for the aroma and flavor of many foods (Tieman et al., 2007); and different bifunctional dihydroflavonol 4-reductases (DFR) (see Supplementary Table 3). DFR catalyzes the first step in the conversion of dihydroflavonols to anthocyanins and are responsible for the production of colored anthocyanins (Boss and Davies, 2001; Davies et al., 2003). The same miRNA candidate was described in the grape miRNA atlas (Kullan et al., 2015) also predicted to target several genes of DFR-like and one CCR.

As for known miRNAs, several members of the miR395 family are differentially expressed at 19 °Brix and at harvest in Bolgheri and in both Bolgheri and Riccione, respectively, when comparing the two cultivars. Moreover, miR395f is differentially expressed also in CS at harvest between Montalcino and Bolgheri. This miRNA has been shown to target genes involved in Sulphate assimilation and metabolism (Liang and Yu, 2010; Kawashima et al., 2011; Matthewman et al., 2012), and hence it could be connected to flavonoid and stilbene pathways as suggested by Tavares et al. (2013).

miR399 family members are also differentially expressed in several comparisons: at 19 °Brix between Riccione and Bolgheri in CS and between Riccione and Montalcino in SG, plus in Montalcino between CS and SG. At harvest, miR399 are differentially expressed in SG in all the three comparisons among vineyards and in Riccione between CS and SG. miR399 is implicated in Phosphate homeostasis being rapidly up-regulated upon Pi starvation (Fujii et al., 2005). miR399 regulatory network has been shown to be important in flowering time (Kim et al., 2011) and was identified as a temperature-sensitive miRNA (Lee et al., 2010), however its characterization in fruit ripening is lacking, although intriguing.

miR396 family members are known to be regulated during organ development, targeting Growth Regulating Factors (Liu et al., 2009; Wang et al., 2011) and also in berry development (Kullan et al., 2015; Cao et al., 2016), and we observed their modulation during berry ripening in our data as well, but more interestingly, they are also differentially expressed between CS and SG in berries sampled in Bolgheri at 19 °Brix.

Finally, the investigation of the global relationships of different small RNA classes and miRNAs expressed in different grapevine cultivars, collected in different vineyards and developmental stages, suggests that although the vineyard may influence their profile of abundance it probably does in less proportion than developmental stage and cultivar. Somehow, this behavior would be expected because although the epigenetic state is dynamic and responsive to both developmental and environmental signals, small RNAs in general and even more miRNAs are well known to play numerous crucial roles at each major stage of plants development (Jones-Rhoades et al., 2006; Chen, 2009, 2012). The results here described are in agreement with those reported in the grapevine miRNA atlas (Kullan et al., 2015), especially with respect to the clustering of berries according to their developmental stage, sustaining the idea that miRNAs influence organ identity and clearly separate green and ripened berries. Also, in the study of the grapevine transcriptome performed by Dal Santo et al. (2013), they observed that other factors such as year and developmental stage had more influence on the gene expression, rather than the environment.

AUTHOR CONTRIBUTIONS

DPP prepared small RNA libraries, performed the *in silico* analysis and wrote the paper. LB conceived the experimental plan and sampled biological material. SDS prepared plant material for RNA extraction, read critically the paper. GDL prepared plant material for RNA extraction, sampled the biological material, read critically the paper. MP conceived the work. MEP supported the lab work, contributed to data analysis and read critically the paper. BM gave a substantial contribution to *in silico* analysis. EM wrote the paper, prepared plant material for RNA extraction, supported small RNA libraries preparation and helped data analysis.

ACKNOWLEDGMENTS

This work was supported by the Doctoral School in Agrobiosciences of Scuola Superiore Sant'Anna and by the Valorizzazione dei Principali Vitigni Autoctoni Italiani e dei loro Terroir (Vigneto) project funded by the Italian Ministry of Agricultural and Forestry Policies. This work benefited from the networking activities within the European funded COST ACTION FA1106 "An integrated systems approach to determine the developmental mechanisms influencing fleshy fruit quality in tomato and grapevine." SDS was financed by the Italian Ministry of University and Research FIRB RBFR13GHC5 project "The Epigenomic Plasticity of Grapevine in Genotype per Environment Interactions". Research in the Meyers lab is supported by the US National Science Foundation Plant Genome Research Program (award #1339229). We wish to thank Jayakumar Belli Kullan for assistance with small RNA libraries preparation, Tzuu-fen Lee for useful discussion on clustering analyses and Mayumi Nakano for curating the database and GEO data submission.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016. 01459

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